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# Embryology of the Rhesus Monkey\*

# Andrew G. Hendrickx and Roger H. Sawyer

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sively in medical research for the past 70 years, embryological studies Although the rhesus monkey (Macaca mulatta) has been used extenhave received little attention. Prior to the report of Heuser and Streeter \*Unpublished data in this Chapter was supported by USPHS research grants RR00169 and HDNS08658 through the California Primate Research Center and School of Medicine, University of California, Davis.

directed toward defining the critical events during organogenesis as a preliminary step to experimental studies (Steffek et al., 1968; Hendrickx, of drugs which pose or may pose some threat to human pregnancies is (1941), which emphasized embryonic development during the first 3 weeks of pregnancy, moststudies were concerned with events associated with ovulation or placentation (Hartman, 1932, 1933; Lewis and Hartman, 1933; Wislocki and Streeter, 1938). More recent studies have been 1972a,b). The value of the rhesus monkey in the teratological evaluation well established (Wilson and Gavan, 1967; Wilson et al., 1970; Wilson, 1972), although it should be recognized that the rhesus monkey is only one of eight species which are potentially useful in this field.

The purpose of this chapter is to present the available information on embryonic development for the rhesus monkey as well as some of the major developmental phases of placentation and fetal growth in order to observations and that much more research is necessary to understand how normal development proceeds and how it may be altered by provide a general account of prenatal development. The reader should bear in mind that in many instances the statements are based on limited environmental factors.

### I. FERTILIZATION

Mastroianni and Brackett (1972) have recently presented the data Fertilization, the fusion of male and female gametes forming a single cell, the zygote or embryo, begins with the entry of the sperm into the outer layer of cells, known as the cumulus oophorus, are dispersed after dispersing the peripheral cells surrounding the ovum has not as yet ing of hyaluronidase and a trypsinlike enzyme which is believed to be cida entering the perivitelline space (Stambaugh and Buckley, 1970). The rypsinlike acrosomal enzyme is located at a subcellular level in ejaculated rhesus monkey sperm and has been shown to exist in an active form prior to exposure of the spermatozoon to the female reproductive which are known about fertilization and related events in primates. cytoplasm of the ovum and ends with the formation of the metaphase plate of the first cleavage division. The ovum is surrounded by cells from the ovary. The cells are organized into two layers; the loosely arranged the ovum enters the oviduct, while the more densely arranged corona radiata layer remains intact. The role of enzymes or spermatozoa in been explored; however, the acrosomal portion of the rhesus monkey spermatozoon has been shown to contain an enzyme complex consistresponsible for the ability of spermatozoon to penetrate the zona pellutract (Stambaugh and Buckley, 1972). From circumstantial evidence, it

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is known that fertilization takes place in the oviduct, but it is not kno in what portion it occurs.

small despite the millions ejaculated into the vagina at copulativ reached the oviduct following insemination directly into the uterus a after natural mating. This was done by counting the number of spe recovered in oviductal washings at specific times after inseminativ evidence suggests that rhesus monkey spermatozoa require capaci tion before they are capable of fertilizing an ovum (Dukelow a the spermatozoon penetrates the ovum, the latter finishes its seco Marston and Kelly (1968) estimated that less than fifty spermatoz Although there is no direct evidence that capacitation occurs, indir Chernoff, 1969). Based on the time of sperm penetration, spermatoz maturation division and both the female and male pronuclei are form The number of sperm which are transported to the oviduct is relativ require 3 to 4 hours for capacitation (Marston and Kelly, 1968). Af The fertile life of ova and spermatozoa is unknown.

like many other mammalian forms (Marston and Kelly, 1968). Once spermatozoon has penetrated the zona pellucida and vitelline me brane and lies within the vitellus, the tail drops off, and the head swe is extruded from the ovum soon after the spermatozoon enters, a Although the exact process by which the sperm penetrates the vii line (yolk) membrane has not been determined, it has been shown t no longer recognizable. Presumably, the pronuclei enter cleave division. O'Rahilly (1973) gives a more complete account of the chang the midpiece and tail are carried with the head into the ovum, mu forming the "male pronucleus" (Fig. 1). The second polar body proba formation of the "female pronucleus" begins. Numerous nucled that occur in the early stages of human development. The same crite are probably applicable to the study of rhesus monkey devele respective pronuclear membranes disappear, and the pronuclei are present in the pronuclei. In the next step of development,

was believed to have occurred some hours before actual mating, a Marston and Kelly (1968) recovered pronuclear stages within 6 ho after insemination, and Lewis and Hartman (1941) recovered a 2-c ovum from the oviduct 23 hours after mating. In the latter case, ovulati it was considered that the ovum may have been in the oviduct prior fertilization and was in an "overripe" condition. This may account for slightly abnormal appearance and its failure to develop further in in vi The time intervals between ovulation and fertilization, and fertili tion to the first cleavage division have not been thoroughly studied, I

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### II. CLEAVAGE

The fertilized ovum or embryo remains in the oxiduct for 3 or 4 days before transfer to the uterus (Marston and Kelly, 1968; Lewis and Hartman, 1933, 1941). The timing of transfer from the oviducts to the uterus and the role of the utero-tubal junction in governing this transfer is not understood, although its importance has been demonstrated in several mammalian species. The normal course of cleavage has not been studied extensively in primates. No evidence is available for the first cleavage division, but the second cleavage division occurs at right angles to the first, the third approximately at right angles to the second. Cleavage divisions are not synchronized, so that both odd- and evennumbered cell stages may be found (Fig. 2).

### A. Cleavage Rates

Information on cleavage rates is scanty indeed; only approximate estimates are available for rhesus monkey embryos (Lewis and Hartman, 1933, 1941). These estimates are based on the detection of ovulation by rectal bimanual palpation of the ovary at various intervals (Hartman, tion and the in vitro culture of the recovered ova. The cleavage stages were estimated as follows: 1-cell stage, 0-24 hours; 2-cell stage, 24-36 1932, 1933) to confirm the duration of early cleavage stages from ovulahours; 3- and 4-cell stages, 36-48 hours; 5-and 8-cell stages, 48-72 hours, and 9- and 16-cell stages, 72-96 hours after ovulation. Information on the transport of embryos down the oviduct is vague, but according to the The developmental stage at which the embryo reaches the uterus may be related to differences in either the rate of cleavage or in the rate of studies cited above, the embryo enters the uterus at the 16-cell stage. tubal transport.

#### B. Blastocyst

enlarges, lining the zona pellucida with a layer of cells in the process. The After the cleaving embryo (morula) enters the uterine cavity, it transforms into a blastocyst. The morula forms a cavity, the blastocoel, that polarity of the embryo is established at this time by the gathering of certain cells at one pole. These cells form the inner cell mass, or embryoblast, from which the embryo develops. The layer of epithelial cells on the inside of the zona pellucida comprises the trophoblast. The morula is converted into a blastocyst 5-6 days after ovulation and subsequent fertilization (Heuser and Streeter, 1941). The zona pellucida is shed 6 or 7 days after ovulation and prior to implantation (Fig. 3).

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## III. IMPLANTATION

Implantation occurs during the blastocyst stage of development a the zona pellucida is shed and the trophoblast comes in contact v the endometrium. Implantation in the rhesus monkey, as in ot primates, involves a sequence of integrated steps which have be Considerable change is seen in the cells comprising the various o reviewed recently by Blandau (1972). The blastocyst remains free in uterine lumen for 9 days before implantation occurs. The size and sh usually resembles a spherical mass. Some differentiation is occurrin into the polar trophoblast (i.e., in the region of the inner cell m which may become multilayered very early, and the single laye trophoblast cells which forms the continuous lining of the blastoc The inner cell mass gives rise to two types of cells—large, lightly stair the free blastocyst at days 8 and 9. The trophoblast is differentia cells (epiblast), and flattened epithelial cells that have delamina stituents in the 8- and 9-day blastocyst, as shown in the tabulation be of the free blastocyst varies considerably (Heuser and Streeter, 1941), from the inner cell mass and constitute the endoderm (endobl

8-Day blastocyst	
Polar trophoblast ceils	. 56
Cavity wall trophoblast cells	28
Epiblast cells (embryonic)	4
Endodermal epithelial cells	12
Total cells	140
9-Day blastocyst	
Polar trophoblast cells	95
Cavity wall trophoblast cells	224
Epiblast cells (embryonic)	32
Endodermal epithelial cells	24
Total cells	375

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it is through the multifaceted functions of this membrane tha The trophoblast of the blastocyst is of special significance bec embryo remains viable and is secured to the maternal endomet (Blandau, 1972). The functions that have been ascribed to it incl (1) developing an attachment cone that initially anchors the blastc to the maternal endometrium; (2) acting as a selective membra control materials entering the blastocoel; (3) assisting in the esca the embryo from the zona pellucida; (4) transforming into the syn trophoblast at the appropriate time in development, and throu adhesiveness and cytolytic capabilities invading the endometrial str

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and (5) producing and secreting both protein and steroid hormones and developing into a complex endocrine organ, the placenta.

The temporal relationship between escape of the blastocyst and its initial attachment to the endometrium for implantation has not been established for primates.

Precise timing and an intricate interplay of hormones on the endometrium are essential for successful embryonic attachment and implantation. This is even more critical in species, including the rhesus monkey and other primates, in which there is no delayed implantation. Some information is available for the hormonal requirements of implantation in the rhesus monkey. Meyer et al. (1968) demonstrated that implantation occurs in the presence of progesterone alone and in the absence of ovarian estrogen. Although ovarian estrogen is not required for implantation, the possibility that the adrenal cortex provides sufficient estrogen for implantation or the possibility that the uterus may have become sensitized by the ovarian estrogen at a stage early looked.

With regards to the orientation of the blastocyst to the endometrium, the trophoblast adjacent to the inner cell mass is the first site of attachment (Figs. 4 and 5). Attachment is preceded by rapid proliferation of the trophoblast cells, as indicated by the difference in numbers of cells in 8- and 9-day blastocysts, and the formation of a syncytium in the trophoblast above the inner cell mass. Alterations in the endometrial epithelium are visible at very early stages of attachment of the blastocyst. The nuclei are deranged and the cytoplasm stains lightly, possibly indicative of beginning erosion and cytolysiš (Blandau, 1972). According to Heuser and Streeter (1941), the trophoblast is probably active in the lysis of the uterine epithelium. During the initial attachment, the uter-Invasion of the uterine stroma is accomplished on the tenth day of development, and the gap filled by the invading embryo is filled by the rapidly developing syncytium. The embryo collapses somewhat as it invades the endometrium. The surface epithelium and the necks of the glands proliferate on the outer boundaries of the embryo. There is no ine epithelium immediately above the inner cell mass is disrupted. evidence of a decidual response in the uterine wall at day 10, or is there an indication of when definitive decidualization begins (Heuser and Streeter, 1941). The attachment or implantation of the rhesus monkey embryo is superficial, attaching itself to the endometrium and remaining partially within the uterine cavity. This is in contrast to the chimpanzee and human embryo, which are completely interstitial in their implant-

## IV. BILAMINAR DISC

Differentiation of the inner cell mass begins as early as day 9, at w time the inner cell mass and trophoblast are indistinct. By the tenth the inner cell mass is actively dividing, and the cells of the epiblas distinguishable from the angiogenic cells. The epiblast consists, firrirregularly arranged cuboidal cells which become pseudostrat columnar by day 11. It increases in size, but otherwise changes little about day 16, when the primitive streak first appears (Figs. 6 and 7). Sendoderm cells are delaminated from the inner cell mass as the blicyst is implanting; however, active proliferation occurs after attach on days 10 and 11. From a few scattered cells the endoderm forms a definite layer beneath the epiblast. The cells of the endoderm forms are loosely arranged and cuboidal in shape, and do not becorganized into a plate until about day 16, at which time the polarity cembryo is established with the formation of the prochordal plathe cranial end, and the development of the body stalk at the cities.

### V. TRILAMINAR DISC

between days 12 and 14 (Luckett, 1971). The caudal portion c with the primitive endoderm to form the gut endoderm. The r primitive streak, which is quite inconspicuous at this stage, is consic to be the primary source of the extraembryonic mesoderm, althoug margins and endodermal cells ventrally. The endodermal cells The caudal portion of the primitive streak develops very precocic trophoblast may provide an additional contribution. The prir streak is readily observed in 16- and 17-day-old embryos (Fig. 7 a It appears as a proliferated area along the craniocaudal axis in the c region of the ectodermal plate. There are localized areas of diso ized cells along its entire length until the last segment is laid dowr dermal cells cluster in their craniolateral movement and form the of Heuser, and initiate the development of the notochord. A blastc depression (primitive pit) forms in the older specimens as the ecto primitive streak gives off embryonic mesoderm along its craniol is elevated by the increased proliferation of cells.

The notochord appears by the nineteenth day as a short, mode thick column of cells extending cranially from the primitive strethe prochordal plate. It is quickly organized into a rodlike mass remains closely opposed to the endoderm and to the neural ectoderm at its cranial end. The notchordal canal, which forms a sectoderm at its cranial end.

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communication between the amniotic cavity dorsally and the vitelline cavity ventrally, appears shortly after the notochord is formed and extends over its full length by the time the somites appear.

The prochordal plate, which along with the body stalk establishes the mark until somite formation is underway. It marks the location of the polarity of the embryo, appears early and remains as a distinct landbuccopharyngeal membrane.

### VI. ORGANOGENESIS

beyond establishing criteria for staging, there is little reason to believe in Figs. 1 through 23. Because the embryo develops at variable rates as exclusive, but rather as modal. The timing of gestation has been tion in a practical way. The gestational ages given in Figs. 1-23 are Studies concerning the development of organs and organ systems in the rhesus monkey are limited to the gonads (van Wagenen and Simpson, 1965) and the palate (Asling and van Wagenen, 1967; Steffek et al., 1968). Although other organ systems have not been studied they differ significantly in their development from other mammalian forms. The main structural changes during organogenesis are outlined after fertilization, the stated days of gestational age must not be regarded reasonably accurate, because of the ability to identify the time of ovulabased on the actual palpation of the ruptured follicle following ovulation (Hartman, 1933), or by matings of only 2hours duration (Hendrickx, 1972b). Staging of embryos by correlating external and internal form with age and size provides the criteria for determining the age of an embryo with an unknown history or calculating the degree of develop-(1972) have been included. The modifications include the designation ment at a particular stage in pregnancy for experimental purposes. Staging follows that originally proposed by Streeter (1942, 1945, 1948, 1951) for human embryos, but modifications presented by O'Rahilly "stage" as a replacement for "horizon," and Arabic numerals have replaced Roman numerals. Figures 1 to 23 are drawings of photographs of actual embryos which represent stages 1-23.

# VII. COMPARISON TO THE HUMAN EMBRYO

Sufficient data are available to compare rhesus monkey and human embryos with respect to developmental stage, crown-rump length, and ovulation age (Fig. 24). Figure 24A shows that the mean line for human embryos is inclined considerably more horizontally compared with that compiled for the rhesus monkey. This indicates that the mean length for

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mental stage in relation to crown-rump length in human and rhesus monkey embryos (Fig. 24B). A comparison of developmental stage with estimated ovulation age shows that human embryos are about 5 days older than rhesus monkey embryos for each developmental stage (Fig. the estimated ovulation age is greater in human embryos at early ages but falls behind at later ages. There is great similarity in the develop-

# VIII. EXTRAEMBRYONIC MEMBRANES

shows the difference in extraembryonic membrane relationships The mode of development of the extraembryonic membranes has only been studied to a limited degree in the rhesus monkey. Figure 25 between Galago and higher forms such as the rhesus monkey.

#### A. Amnion

exocoelomic space, and completes its reflections about the embryo cellular spaces between the amnion cells and the embryonic disc. The they differentiate from the cytotrophoblast. By day 12, delamination of is laid down between them. Extraembryonic mesoderm occupies the From this point, the amnion expands (Figs. 6 and 7) at the expense of the and umbilical cord. In some instances, a secondary connection between the amnion and chorion exists, resulting in the formation of the amniotic duct passing from the caudal extremity of the amnion to open onto the surface of the trophoblast. Although it is clearly a secondary structure resulting from caudal prolongation of the amnion along the Amniogenesis begins on day 10 of development (Heuser and Streeter, 1941). At this time, a layer of amniogenic cells is clearly demarcated from the embryonic disc and is in the process of being delaminated from the trophoblastic wall. At the same time, fluid accumulates in the interamniogenic cells become aligned into a single-layered membrane as the amnion from the trophoblast is advanced, and a reticular network reticular space, and by day 13 it forms the second layer of the amnion. developing body stalk, it is common to many Old-World monkeys, apes, and man (Hill, 1932).

# B. Allantois and Body Stalk

There are few details available on development of the allantois in the rhesus monkey; it remains rudimentary and plays no major role in 151

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development. The allanto-enteric diverticulum quickly subdivides into allantois and cloaca, and the allantois extends about one-half the way into the body stalk. Its endodermal portion is slightly thicker than the lining of the vitelline sac.

The body stalk develops prior to the appearance of the primitive streak. It is sometimes described as having two parts—a short, cylindrical, distal part, continuous with the chorionic mesenchyme, and a longer, proximal part, attached to the amniotic ectoderm. The latter part forms the posterodorsal wall of the amniotic cavity. The body stalk consists first of extraembryonic mesoderm and is later reinforced by mesoderm from the primitive streak. By the time the somites begin to form, the mesenchyme has condensed to form a solid stalk containing umbilical vessels, and by the time somite formation is complete, the umbilical cord (body stalk) has rotated into a ventral position, the embryo is coiled around it, and the amnion sheathesits proximal surface. Angiogenesis probably occurs simultaneously in the yolk sac, body stalk, and chorion.

#### C. Yolk Sac

ment, cells delaminate from the cytotrophoblast to form Heuser's Yolk sac development in the rhesus monkey is believed to differ slightly from that in humans. In the human embryo, on day 9 of developmembrane, which is continuous with the edges of the endoderm layer. Together with the endodermal layer they form the primary yolk sac (Langman, 1969). In the rhesus monkey, this membrane is formed by migration of the primary endoderm cells along the inside of the blastocyst cavity (Heuser and Streeter, 1941). Extraembryonic mesoderm is laid down between the trophoblast and primary yolk sac by day 12 in both species and, at about this time, endoderm cells begin to spread over the inside of Heuser's membrane. This process continues until the evidence that in both species, large portions of the definitive yolk sac are pinched off (Hertig et al., 1956). The possibility exists that the definitive yolk sac develops by opening of the endodermal plate and the newly formed cells gradually line a new cavity, the definitive yolk sac. The definitive yolk sac is smaller than the primary yolk sac, and there is primitive yolk sac is pinched off in toto and, as a result, plays no role in the formation of the definitive yolk sac (Strauss, 1945; Starck, 1956). The yolk sac remains small until about day 17, when distension begins. Although it is regarded as a vestigial structure, it is not uncommon to see the vesicle on the surface of the placenta at term.

### IX. PLACENTATION

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### A. Formation

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Of the extraembryonic membranes, the chorion is exclusively involved in the formation of the placenta, except in prosimians, such as Calago sp., in which the allantois also plays a role in placentation. As the placenta forms, the chorion takes on two forms—the membranous chorion, which extends between the two placental discs, and the villous chorion which is anchored to the endometrium. The rhesus monkey, like the majority of New- and Old-World monkeys, has a bidiscoid placenta which consists of two separate placental discs—a primary disc, the first site of attachment at the embryonic pole, and a secondary disc at the abembryonic pole.

Development of the placenta can be divided into three stages (Wislocki and Streeter, 1938):

# 1. THE PRELACUNAR STAGE (PERIOD OF TROPHOBLASTIC PLATE)

This stage encompasses the ninth and tenth day of development and is characterized by the presence of a solid trophoblastic plate at the primary implantation site which invades and destroys the uterine epithelium at the point of blastocyst attachment. The maternal reaction is proliferation of the uterine epithelium and the uterine glands. In the older embryos of this stage, the only sign of a secondary placenta at the abembryonic pole is some slightly enlarged trophoblastic cells. Proliferlation of the endometrial epithelium and the cells along the necks of the glands at the secondary site is equal to that seen at the primary site.

# 2. THE STAGE OF TROPHOBLASTIC LACUNAE

This stage extends from approximately the eleventh to the fifteenth day. The lacunae, formed by the differentiation of the trophoblastic plate into a reticulated mesh consisting of syncytial trophoblast, fill with maternal blood and proliferation of the uterine epithelium increases. In 11-day-old embryos, the trophoblast at the secondary implantation site begins to proliferate but does not attach to the already proliferated endometrial epithelium. By the 12th day of development, actual attachment occurs and invasion of the trophoblast into the endometrium begins (Fig. 26A,B).

### 3. THE VILLOUS STAGE

This stage extends from about the 15th to the 35th day and is characterized by the formation of the chorionic villi. The villi form by the

differentiation of the cytotrophoblastic cell columns and the transformawhich form the villi cores. Proliferation of the uterine epithelium ceases tion of part of the cytotrophoblast into mesenchyme and angioblasts at the beginning of this period. The uterine epithelium is replaced at the fetal maternal border by the junctional zone. The fetal part of the border is converted into a typical trophoblastic shell by the fusion of the expanded ends of the cytotrophoblastic cell columns into a distinct plate (Fig. 26C,D). The formation of the villi in the secondary placenta follows the same pattern as in the primary placenta but is delayed by several

# 4. DEFINITIVE PLACENTA

capillaries throughout gestation, but certain maternal tissues are broken of membranes or cell layers comprising the placental barrier. In the directly. At the ultrastructural level there are further differences in the ing the rhesus monkey, consists of one layer, thus, it is classified as the hemomonochorial type (Luckett, 1970; Panigel, 1970), in contrast to the hemotrichorial type found in several laboratory species. A comparison Formation of the definitive placenta is essentially complete by day 35 of development (Fig. 27C). Minor changes occur, including: (a) intervillous connections from strands of syncytium, (b) increased branching and refinement of the villi with age, and (c) a reduction of the chorionic vessels on the placental surface and those connecting the secondary placenta to the primary placenta, so that a number of vessels connect the from the chorion is the primary means of increasing the surface of contact between maternal and fetal tissues. The distribution of villi is imited to two discs, the primary and secondary, hence, the term bidiscoid placenta (Fig. 27D). Within the disc, the villi are extremely long and have a complex branching pattern. The chorion remains intact over the fetal down and disappear during placental formation and reduce the number placenta of the rhesus monkey, the endometrial epithelium, the endometrial stroma; and the maternal capillary endothelium are all destroyed and maternal blood bathes the chorion (trophoblastic surface of villi) trophoblast (Enders, 1965). The trophoblast of higher primates, includof the specific structural differences in the rhesus monkey, baboon, and two discoidal placental masses. The outgrowth of the chorionic villi man is shown in Table I.

### B. Circulation

pective blood streams are brought into close proximity so that an efficient The passage of materials between the mother and embryo (fetus) occurs within the placenta. In the process of development, their res-

Embryology

TABLE 1 Comparison of the Main Placental Features of the Rhesus Monkey, Man, and Baboon\*

The same of the sa		
Rhesus monkey	Man	Baboon
Implantation Superficial No decidual reaction	Interstitial Pronounced decidual reaction	Superficial (central) Partial decidual reaction; developing somewhat more slowly than in man
Transitory epithelial plaque No epithelial reaction Secondary placenta  No secondary placent	No epithelial reaction No secondary placenta	No epithelial reaction No secondary placenta
Trophoblast Boundary between maternal and fetal	Boundary between maternal and fetal	Boundary between mater- nal and fetal tissues—
tissues—straight No penetration of myometrium by	tissues—very irregular Penetration of inner third of myometrium by	straight  No penetration of myo- metrium by trophoblast
trophoblast No trophoblastic wandering cells	trophoblast Trophoblastic wandering cells prominent	No trophoblastic wander- ing cells
Arteries Intravascular cells very early—17th day Migration of intravascular cells in lumen only Elastic tissue extending half-way up in endometrium Multiple openings to intervillous space from single stems— inframment	Intravascular cells later— peak at 12th week Walls also traversed by intravascular cells Little elastic tissue beyond myoendometrial junction Multiple openings— common	Intravascular cells very early—16th day Intravascular cells in lumen only Elastic tissue extends well into endometrium Multiple openings—common

<sup>&</sup>lt;sup>a</sup>From Houston (1969). Amer. J. Anat. 126, 1-15.

villi which contain the blood vessels (Ramsey and Harris, 1966; Harris The first column represents the structure of the villous tree, and column two demonstrates the fetal circulation and how blood vessels pass through the villi. The first two columns further demonstrate that the in the placenta is dependent upon the configuration of the chorionic and Ramsey, 1966). A composite drawing showing the structure and corresponding circulation of a hemochorial placenta is shown in Fig. 28. villus is firmly attached to the basal plate with free-floating branches protruding into the intervillous space along the length of the mair villous trunk. Each villous trunk carries both arterial vessels, which branch and extend into the smaller terminal villi, and venous vessels which return the blood through the same villous trunk. There are no interchange may take place. The structure of the fetal circulation with-

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anastomoses between villi, or between the vascular channels within a villus.

ation of the maternal blood. In the discoid, hemochorial placenta, there pressure in the maternal vessels entering at the base of the placenta is The remaining three columns of Fig. 28 represent the placental circulwithin the intervillous space is not strictly random. Because the blood much greater than in the intervillous space, the blood is driven toward the chorionic plate, as shown in column four. The loose, semiattached fronds of villi which are encountered along the course of the blood stream baffle the flow and direct some of the blood laterally. The force of the blood entering the intervillous space pushes blood out through in the absence of preformed channels, the flow of maternal blood the circulatory pathway of the maternal blood is governed by the pressure it is under when it enters the intervillous space. The spurting pattern the venous channels, which are also located in the basal plate. As a result, of maternal blood into the intervillous space varies with the contracted out pregnancy and vary greatly in intensity. The degree of contraction are no preformed channels for the passage of maternal blood; instead the maternal blood enters at the base into the space between the villi or relaxed state of the uterus. Contractions of the uterus occur throughgoverns the rate of flow.

### X. FETAL PERIOD

In human embryology it is customary to refer to the conceptus as a fetus toward the end of the eighth week postconception, when major organogenesis is essentially completed. This custom is gradually becoming accepted in nonhuman primate embryology as well, and for the rhesus monkey the conceptus is referred to as a fetus by about day 45. The most conspicuous developmental process during the fetal period is physical growth. By the process of histogenesis, the individual organs, established as simple structures during organogenesis, are converted into more specialized ones with a primary function. Functional matur-1974), although it is of major importance. Environmental factors which influence physical growth are the availability of nutrients through the maternal diet and placental transport. The endocrine balance and the absence of disease also contribute to the maternal physiological state but work has been done (Kerr et al., 1969) to provide sufficient data ation of individual organs has received only limited attention (Rakic, on normal organ growth of the rhesus monkey fetus (Tables II, III, and Studies related to physical growth of the fetus are somewhat limited, and are of considerable importance to the normal growth of the fetus.

TABLE II The Growth Rates of Fetal Organs of the Rhesus Monkey<sup>2,5</sup>

,	50-75	75-100	100-125	100-125 125-150 150-175	150-175
Organ	Days	Days	Days	Days	Days
Lungs	503.67	82.13	24.60	18.77	-13.60
Heart	317.65	101.55	46.10	21.76	13.49
Liver	447.53	89.82	23.93	23.46	2.80
Adrenals	318.72	. 44.57	15.51	14.04	60.02
Kidneys	572.13	139.74	26.97	9.37	10.04
Thyroid	I	105.52	31.23	32.72	20.07
Thymus	ı	115.54	59.69	24.18	-20.92
Spleen	119.24	120.07	33.04	19.74	6.64
Brain	ł	69.66	49.87	10.28	4.74
Placenta	73.02	24.86	7.21	17.79	9.77
Fetus	392.40	98.01	39.12	23.31	6.59

<sup>a</sup>Derived from change in mean weight of organs at each gestational age. Figures indicate value ±1 SD. Data in mg/gm/day.

<sup>b</sup>From Kerr et al. (1969). Growth 33, 201-213.

TABLE III The Growth in Weight of Major Organs during Fetal Life of the Rhesus Monkey<sup>2,5</sup>

,			Gestational age (days)	age (days)		
- Organ	20	75	100	125	150	175
Total body wt.	4.003	43.27	149.29	295.30	467.38	544.4
	+0.987	±3.49	+9.73	±27.00	±42.76	±101.6
Placenta	$\bar{2}_{1.23}$	59.99	77.76	114.82	165.88	206.40
	+6.59	±12.23	±14.66	±21.28	±33.82	±49.52
Brain	1	5.26	18.36	41.25	51.85	58.00
		+0.64	±1.15	±3.57	±3.04	±5.97
Spleen	0.017	0.068	0.271	1.495	0.739	0.861
		+0.014	±0.071	0.09€	±0.131	±0.265
Thyroid	I	0.015	0.054	0.096	0.175	0.263
		+0.007	±0.011	±0.052	090.0∓	$\pm 0.137$
Kidnevs	0.017	0.225	1.145	1.92	2.37	2.96
	+0.007	080.0∓	+0.155	±0.37	±0.37	<b>∓0.67</b>
Adrenals	0.007	0.065	0.138	0.192	0.259	0.649
	+0.003	+0.022	±0.026	±0.026	±0.100	±0.279
Liver	0.148	1.80	5.84	9.33	14.81	15.85
	+0.061	±0.40	€9.0∓	±1.38	±1.49	±2.84
Heart	0.025	0.226	0.800	1.72	2.66	3.55
	+0.006	±0.048	060.0∓	±0.39	<b>±0.3</b> 5	±1.22
Lunes	0.097	1.32	4.01	6.48	9.53	6.29
Ь	+0.041	+0.20	+0.88	±1.42	±1.27	+0.99
Thymus	ļ	0.086	0.333	0.829	1.331	0.635
		±0.35	±0.126	±0.277	±0.432	±0.223

<sup>a</sup>Figures indicate mean weight  $\pm 1$  SD. All data in gm. <sup>b</sup>From Kerr et al. (1969). Growth 33, 201–213.

TABLE IV The Relative Growth Rate of Fetal Organs of the Rhesus Monkey<sup>a,b</sup>

		0.0000	1 1	1		
		Cestatio	Gestational age (days)	(5)		
Organ	20	75	100	125	150	175
Placenta	551.7	140.6	65.2	38.8	35.8	34.4
	±201.5	±29.9	+9.1	<del>1</del> 5.9	+8.1	+3,3
Brain	ı	12.14	12.30	14.02	11.15	11.14
		+0.99	±0.49	±1.18	+0.88	±1.88
Spleen	0.70	0.16	0.18	0.17	0.16	1.6
		±0.03	₹0.05	+0.03	±0.02	+0.04
Thyroid	1	0.034	0.036	0.033	0.037	0.047
		+0.016	±0.008	±0.019	±0.010	+0.020
Kidneys	0.536	0.587	0.765	0.620	0.507	0.548
,	±0.061	±0.164	+0.071	±0.118	±0.038	+0.102
Adrenals	0.233	0.151	0.093	0.064	0.055	0.113
	$\pm 0.039$	±0.055	±0.017	±0.012	±0.021	+0.037
Liver	4.05	4.14	3.90	3.16	3.19	2.93
	±1.27	±0.71	±0.25	±0.31	±0.30	+0.35
Heart	0.657	0.525	0.543	0.581	0.561	0.646
	±0.137	±0.083	±0.030	±0.101	±0.043	+0.147
Lungs	2.63	3.04	2.68	2.19	2.04	1.22
1	±0.47	±0.43	±0.47	±0.42	±0.14	±0.26
Thymus	1	0.201	0.224	0.280	0.283	0.118
		₹0.093	₹0.092	±0.082	+0.080	+0.035

<sup>&</sup>lt;sup>2</sup> Figures indicate mean value ±1 SD. Data given in percentage total body weight. <sup>b</sup>From Kerr et al. (1969). Growth 33, 201–213.

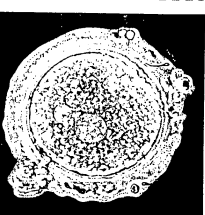
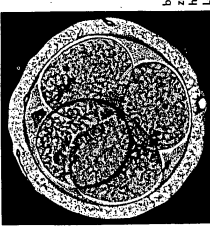


Fig. 1. Stage 1. 0–1.0 days post fertilization. One-celled ootid, male and female pronuclei present, corona radiata dispersed, sperm seen in zona pellucida, probably found in upper half of oviduct (after Suzuki and Mastroianni, 1968).

The state of the

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blastomeres, 2- to 16-cell stages to morula zona pellucida persists, located in lowe half of oviduct and uterine cavity (afte Fig. 2. Stage 2. 2-4 days. Segmenting Lewis and Hartman, 1941).

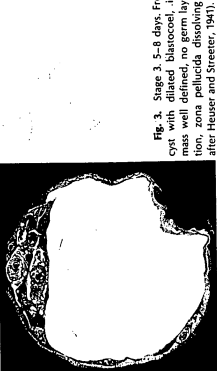


Fig. 3. Stage 3. 5-8 days. Free blastc cyst with dilated blastocoel, inner ce mass well defined, no germ layer form: tion, zona pellucida dissolving (section

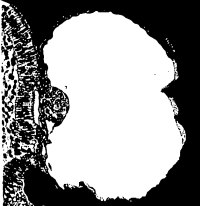


Fig. 4. Stage 4. 8-10 days. Attachir blastocyst, blastocoel distended, trophe blast disrupting epithelium and impingir upon stroma, germ layer formation begin amniogenic cells appear (section, aft Heuser and Streeter, 1941).



blast proliferating at embryonic pole, no villi, no maternal decidual response, amniotic cavity dilates, extraembryonic Fig. 5. Stage 5. 10-11 days. Tropho-(transverse section, after Heuser and Streeter, 1941). mesoderm proliferating



Fig. 6. Stage 6. 12-15 days. Primitive villi appear, yolk sac forms, bilaminar germ disc distinct, primitive streak begins to form, body stalk indicated (transverse section, after Heuser and Streeter, 1941).



Fig. 7. Stage 7. 16-18 days. Branching streak formed, angioblasts appear in the villi appear, axis of embryonic disc deterrnined by head process. Notochordal derm begins to delaminate, primitive yolk sac and chorionic membrane (transprocess appears, intraembryonic mesoverse section, after Heuser and Streeter,

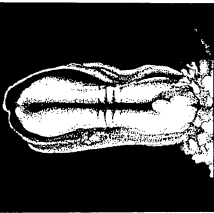
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canal forming, primitive streak well estabneural plate appears (sagittal section, Fig. 8. Stage 8. 19-20 days. Embryonic definite body stalk present, primitive pit and neurenteric lished, intraembryonic mesoderm formed, after Heuser and Streeter, 1941). elongating,



somites. Obvious head fold, caudal fold foregut forms, primitive streak extends from cloacal membrane to neurenteric body stalk elongates (after Heuser and Fig. 9. Stage 9. 20-21 days; 0-3 paired appears, neural folds and goove distinct, canal and occupies 1/3 to 1/4 the length of the embryo, notochord prominent, Streeter, 1941).

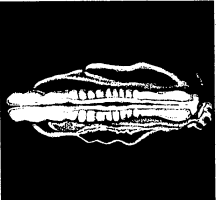


Fig. 10. Stage 10. 21-23 days; 4-12 somites. Neural folds begin fusing to form neural tube, optic primordia form, otic placodes readily detectable, 1st branchial hindgut elongates (after Heuser and arch (mandibular) evident, endocardial tubes fuse, 1st and 2nd aortic arches form, Streeter, 1941).

Fig. 11. Stage 11. 24-26 days; 2-3.5 mm; 13-20 paired somites. Cranial neuropore closes, optic vesicle evaginates, cranial nerves V, VII, VIII, IX, X, and XII appear, S-shaped heart with sinus venosus prominent, mandibular and hyoid arches well defined, 1st and 2nd pharyngeal pouches form, liver primordium forms, yolk stalk begins, body axis elongated and usually curved (after Heuser and Streeter,



12. Stage 12. Z-28 days; 3-5 arches present, 3rd aortic arch appears, oropharyngeal membrane ruptures, optic vesicle contacts surface ectoderm, otic and forelimb bud forming, yolk stalk mm; 21-29 paired somites. Three branchial pit closing, appendicular ridge distinct formed, respiratory diverticulum evaginates, caudal neuropore closes, body axis C-shaped (after Heuser and Streeter,

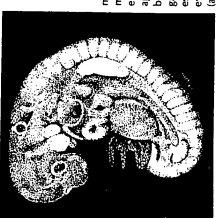
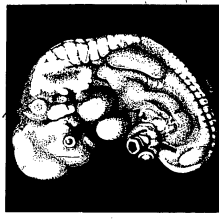


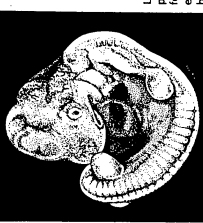
Fig. 13. Stage 13. 28-29 days; 4-6 maxillary process appears, lens placode Four branchial arches present, evident, otocyst closed, olfactory placode appears, Rathke's pouch forms, hindlimb esophagus and trachea divide, stomach evident, mesonephric ducts join cloaca bud appears, thyroid primordium bilobed, gallbladder and dorsal pancreas appear, (after Heuser and Streeter, 1941).

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tiated, lung buds elongate, ventral Fig. 14. Stage 14. 29-30 days; 5-8 mm. Optic cup and lens placode incervical sinus begins, Rathke's pouch prominent, pharyngeal pouches differenvaginate, endolymphatic duct forms, pancreas appears, metanephric bud forms.



elongate and become flipper-shaped, leg pigment appears in retina, olfactory pits Fig. 15. Stage 15. 31-32 days; 6-9 mm. Lens vesicle closes, optic stalk prominent, form, neurohypophysis appears, arm buds buds elongate, cecal swelling appears.

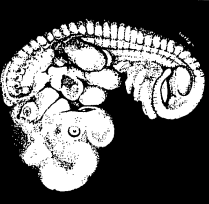


Fig. 16. Stage 16. 33-34 days; 8-11 mm. Pigment in retina prominent, neurohypophysis evaginates, cervical sinus closes, auditory hillocks appear, secondary into umbilical cord, rotation of gut begins, bronchi appear, midgut loop herniated hand plate evident, genital ridge appears, ureteric bud expands into renal pelvis. 5. Embryology

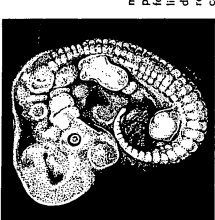
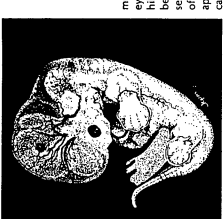
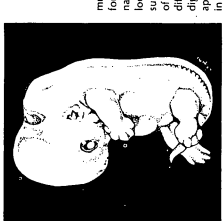


Fig. 17. Stage 17. 35-36 days; 10-12 foot plate appears, tertiary bronchi form, rotates, Müllerian ducts appear, major mm. Nose and primary palate form, hand plate prominent and digital rays appear, dorsal and ventral pancreas fuse, stomach iver enlarges and biliary ducts appear, calices of kidney appear.

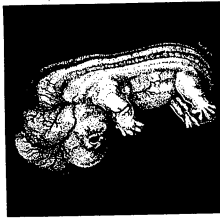


mm. Facial primordia united, eyelids and hillocks forming auricle, neurohypophysis semicircular ducts forming, digital rays of hand prominent, digital rays of foot appear, Müllerian duct prominent, minor Fig. 18. Stage 18. 37-38 days; 12-16 eyes begin to move forward, auricular begins to fold, cochlear duct L-shaped, calices formed, collecting tubules appear.

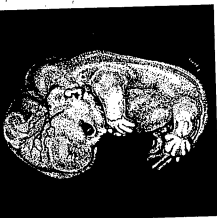


digital notches present, cartilage cells Fig. 19. Stage 19. 38-39 days; 15-19 mm. Semicircular ducts formed, cornea formed, cochlear duct J-shaped, bucconasal membrane ruptures, auricular hillocks coalescing, secondary palate begins, submandibular gland appears, primordium of secretory tubules of kidney form, testis differentiates, forelimb rotating and interappear in humerus, digital rays prominent

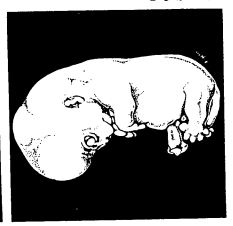
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mandibular gland, ossification centers Fig. 20. Stage 20. 39-40 days; 17-20 mm. Optic nerve fibers reach brain, eyelids cover 1/5 of eye, palatine processes present beside tongue, palms directed caudally, primary branches appear in subindicated by clearing of cartilage, interdigital notches present in foot, anal membrane ruptures.



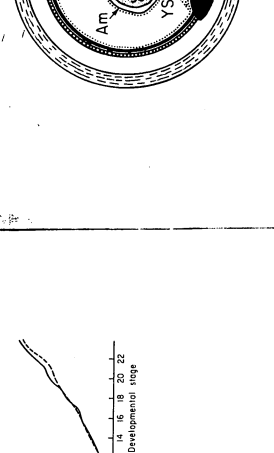
turn, collecting and secretory tubules fuse in kidney, osteoblasts appear in mm. Three layers present in cornea, eyelid covers 1/4 of eye, cochlea completes one Fig. 21. Stage 21. 41-42 days; 19-23 humerus.



process rotated 45° medially, large glomeruli present, calcification begins ir covers 1/3 of eye, secondary branches appear in submandibular gland, palatine Fig. 22. Stage 22. 43-44 days; 22-26 mm. Eye rotated to front of face, eyelid humerus, hindlimb rotating.

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Stage 23. 45-46 days; 25-30 mm. Head raised from chest, eyelids cover most or all of eye, hands overlap in front of face, soles of feet apposed, palatal closure begins, cochlea completes 2½ turns, tertiary branches appear in submandibular gland, midgut hernia withdrawing from umbilical cord, ovary recognizable.



14 16

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Ovulation age (days)

(mm) qmui-nwoi) だめでです

5 2

Crown-rump (mm)

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Fig. 24. (A) Correlation of estimated ovulation age and crown-rump length in rhesus monkey (---) and human embryos (---); (B) Correlation of developmental stage 1968; data for the rhesus monkey are taken from Heuser and Streeter, 1941, and California and crown-rump length in rhesus monkey and human embryos; (C) Correlation of developmental stage and ovulation age (data for man are taken from Nishimura et al., Primate Research Center Collection).

Developmental stage

 $\alpha$ 

found among nonhuman primates. The amnion (Am) remains essentially the same in Fig. 25. Examples of different types of extraembryonic membrane configurations all species, but the allantois (All), which is large in Galago (A) and most other prosimians, is rudimentary in higher primates (B). YS, yolk sac; Ex, exocoelom; BS, body stalk; C, chorionic vesicle; P, placenta. (From Hendrickx and Houston, 1971).

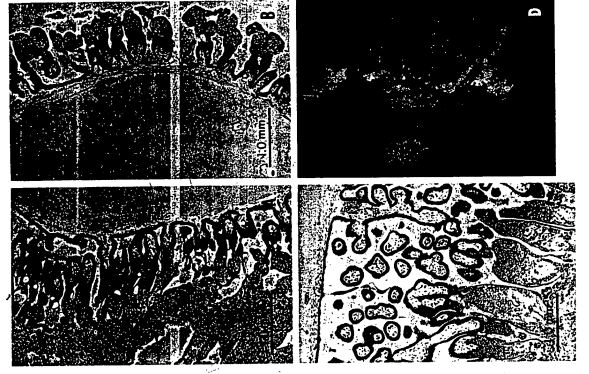


Fig. 26. Development of the placenta. The primary and secondary placental sites are shown for a 14-day-old embryo (A and B) and for a 17-day-old embryo (C and D). (A) Villi have not yet formed at 14 days but there is a broad communication between maternal blood channels and lacunae at the primary site. (B) Proliferative epithelium characterizes the secondary site. (C) Definitive villi with incomplete vascularization separated by lacunae (intervillous spaces) characterize the primary placenta at 17 days. (D) The villi at the secondary site are less advanced, lacking central stroma. A junctional zone, a necrotic area between the maternal and fetal tissues, is evident at both sites (from Wislocki and Streeter, 1938).

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Fig. 28. A composite drawing of the hemochorial placenta (of a man or monkey) to show its structure and circulation. Drawn by Ranice W

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